PATENT ABSTRACTS OF JAPAN

(11)Publication number:

01-135540

(43) Date of publication of application: 29.05.1989

(51)Int.CI.

B01J 29/08 B01D 53/36 B01J 29/18

(21)Application number: 62-292914

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(22) Date of filing:

19.11.1987

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(54) PRODUCTION OF CATALYST FOR PURIFYING EXHAUST GAS

(57) Abstract:

PURPOSE: To improve purification performance of NOx in the lean atmosphere by allowing a monolithic carrier to be spread with slurry in which zeolite, silica sol, alumina sol and water are blended and calcining it and thereafter immersing it in a metallic salt aq. soln.

CONSTITUTION: Slurry is prepared by mixing zeolite, silica sol, alumina sol and water. A monolithic carrier is spread with this slurry and thereafter calcined to form a catalyst carrier. Then a catalyst for purifying exhaust gas is produced by immersing this catalyst carrier in a metallic salt aq. soln. of metal utilized as the catalyst and allowing zeolite to be carried with the metal utilized as the catalyst by ion exchange. A mixing ratio of silica sol and alumina sol is preferably regulated to such mixing ratio that Si/Al thereof is not made far from Si/Al of zeolite.

LEGAL STATUS

[Date of request for examination]

[Date of sending the examiner's decision of rejection]

[Kind of final disposal of application other than the examiner's decision of rejection or application converted registration]

[Date of final disposal for application]

[Patent number]

[Date of registration]

[Number of appeal against examiner's decision of rejection]

[Date of requesting appeal against examiner's decision of

rejection]

[Date of extinction of right]

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File: JPAB May 29, 1989 L8: Entry 8 of 35

DOCUMENT-IDENTIFIER: JP 01135540 A

TITLE: PRODUCTION OF CATALYST FOR PURIFYING EXHAUST GAS

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Abstract (2):

CONSTITUTION: Slurry is prepared by mixing zeolite, silica sol, alumina sol and water. A monolithic carrier is spread with this slurry and thereafter calcined to form a catalyst carrier. Then a catalyst for purifying exhaust gas is produced by immersing this catalyst carrier in a metallic salt aq. soln. of metal utilized as the catalyst and allowing <u>zeolite</u> to be carried with the metal utilized as the catalyst by ion exchange. A mixing ratio of silica sol and alumina sol is preferably regulated to such mixing ratio that Si/Al thereof is not made far from Si/Al of zeolite.

⑫ 公 開 特 許 公 報 (A)

Mint Cl.4

識別記号

庁内黎理番号

❷公開 平成1年(1989)5月29日

平1-135540

29/08 B 01 J 53/36 29/18 B 01 D B 01

A-6750-4G A-8516-4D A-6750-4G 104

未請求 発明の数 1 (全5頁) 審査請求

69発明の名称

排気ガス浄化用触媒の製造方法

創特 昭62-292914

魯田 昭62(1987)11月19日

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1. 発明の名称

排気ガス浄化用触供の製造方法

2.特許請求の範囲

-) 以下の工程:
- (a) セオライト、シリカゾル、アルミナゾル 及び水を混合してスラリーを調製する工程、
- (b) 上記スラリーを一体型担体に付着させ、 焼成する工程、
- (c) 触媒用金属の金属塩水溶液に浸漉してせ オライトに触媒用金属をイオン交換する工

からなることを特徴とする排気ガス浄化用触 媒の製造方法。

- 2) シリカゾルとアミルナゾルの混合比は、そ れらのSi/Ae 比がセオライトのSi/Ae 比 にほど遠くはならない混合比であることを特 徴とする特許請求の範囲第1項記載の方法。
- 3) 工程(c)は、工程(a)及び(b)よりも後で行なわ れる工程であることを特徴とする特許請求の

鉱開第1項記載の方法。

3.発明の詳細な説明

く産業上の利用分野>

本発明は自動車の排気ガス浄化用触媒、特に は空燃比が、リーン側となる酸素過剰雰囲気に おいてもNOxを高率に浄化できる触媒の製造方 法に関するものである。

く従来の技術>

自動車の排気ガス浄化用触媒として、一酸化 炭素(CO)及び炭化水素(HC)の酸化と、窒素 酸化物(NOx)の還元を同時に行う触媒が汎用 されている。とのよりな触媒は、例えば特公昭 58-20307号公報にもみられるように、耐火 性担体上のアルミナコート層に、 Pd. Pt. Rh 等の貴金禺、及び協合により助触媒成分として Ce.La 等の希土類金属又は Ni 等のペースメタ ル酸化物を抵加したものが殆んどである。

かかる触錐は、エンジンの設定空機比によっ て浄化特性が大きく左右され、希梅混合気つま り空燃比が大きいリーン側では燃焼後も酸素(O.)

PTO 2003-3197

の量が多くなり、酸化作用が活発に、還元作用が不活発になる。この逆に、空燃比の小さいリッチ側では酸化作用が不活発に、還元作用が活発になる。この酸化と還元のバランスがとれる理論空燃比(A/F=146)付近で触媒は殷も有効に働らく。

従って、触媒を用いる排気ガス浄化装置を取付けた自動車では、排気系の酸素濃度を検出して、混合気を理論空燃比付近に保つようフィードバック制御が行なわれている。

く発明が解決しようとする問題点>

•

一方、自動車においては低燃役化も要請されており、そのためには通常走行時なるべく酸素 過剰の混合気を燃焼させればよいなとが知りの 砂素過剰雰囲気となって、排気ガス中の有害成 分のうちHC、COは酸化まできても、NOxは 触媒床に吸着したOxによって活性金属との投触 が妨げられるために、強元除去できないという 問題があった。そのため従来、触媒によって高

ナゾルの混合物 50~100 部、水 30~100 部の割合で混合するのが好ましく、 更には pH 調整剤を添加して pH 5~6 のスラリーとするのが良い。

シリカソルとアルミナソルの混合比は、それらのSi/AB比がゼオライトのSi/AB比にほど遠くはならない混合比であることが肝要であり、好ましくは两ソル混合物のSi/AB比がゼオライトのSi/AB比の1/3~5倍となるように、シリカソルとアルミナソルを混合するのがよい。ゼオライトとしては、NOx分子径よりも値かに大きい5~10Å径の細孔を有するゼオライトが適当である。

工程(b) において、上配の一体型担体としては 汎用されているセラミック製モノリス、ハニカム型担体で十分であり、該担体に上記スラリー を付着させるには、担体にスラリーを噴射盤布 するか成はスラリー中に担体を浸漉すればよい。 焼成は、余分な付着スラリーを圧縮空気又は真 空引きにより吹き払ってから、800で以下の巡 度の排気ガス浄化を図る自動車にあっては混合 気を希薄にすることができなかった。

本発明は上記問題点を解決するために為されたものであり、その目的とするところは、リーン側でもNOxを還元除去でき理論空燃比からリーン側の広い領域にわたって全ての有害成分を十分に除去し得る排気ガス浄化用触媒の製造方法を提供することである。

く問題点を解決するための手段>

本発明の排気ガス浄化用触媒の製造方法は、 以下の工程:

- (a) セオライト、シリカゾル、アルミナゾル及 び水を混合してスラリーを調製する工程、
- (b) 上記スラリーを一体型担体に付着させ、焼成する工程、
- (c) 触媒用金属の金属塩水溶液に浸漉してゼオ ライトに触媒用金属をイオン交換する工程 からなることを特徴とする。

工程(a)のスラリーを調製する工程においては、 セオライト100部に対し、シリカゾルとアルミ

度で行なりのがよい。

工程(c)でイオン交換させる触媒用金属としては、Cu, Cr, Co, Ni, Fe, Mn等の選移金属やPt, Pd, Rh, Ce, Ir, Ru等の貴金属が挙げられる。イオン交換は上配金属の水溶性塩りで水溶液中にゼオライトを浸渍するととには001~01mos/sが適当であり、例えば酢酸鍋水溶液の場合、特に002~006mos/sが好ましい(第2図参照)。溶液温度は30±10で充分であり、イオン交換時間は5~140時間、望ましくは40~100時間である(第1図参照)。

本発明の製造方法によれば、工程(a)→(b)→(c)の履でも、また工程(c)→(a)→(b)の顧でも目的とする触媒を得ることができるが、前者の履によるのがイオン交換効果上好ましい。その理由は、イオン交換される金属は交換容易なサイトからより困難なサイトへと段階的に、例えば第3図の構造模式図で示すようにA→B→Cの順に入り、またサイトはC→B→Aと逆順でNOx争化

に有効→無効となる場合を考えれば、最初に全部のゼオライトをイオン交換するよりも第4図に示すように、担体4上のパインダー3間に埋むれたゼオライト粒子2,2 ---- を除いた表面に現われているゼオライト粒子1,1 --- のみをイオン交換したほうが、イオン交換金属5,5 ---- はNOxに対しより高活性なサイトまで入る確率が高くなって触媒活性が向上することになるからである。

く作用>

本発明の製造方法によって得られる触媒は、その表面が、触媒用金属でイオン交換されたゼオライトで優われることとなる。ゼオライトはNOx分子の大きさと並ぶ数点単位の細孔を有しており、そのため該細孔にNOxが選択的に取り込まれる。細孔中には上記金属がイオン交換されて活性となったサイトが存在するため、そこに吸着したNOxは触媒用金属により還元される。
〈実施例〉

以下、実施例により本発明を更に詳しく説明

及透培養機中、70時間室温でイオン交換を 行った。余分を水分を吹き払った後80℃で 20分間乾燥して排気ガス浄化用触媒Aを製 造した。

奥施例 2. 3

実施例1で用いたゼオライトの代わりに、モルデナイト(Si/Ae比190)ならびにフォージャサイト(ゼオライトY; Si/Ae比26)を用い、それぞれのSi/Ae比に合わせたパインダーで調製したスラリーを担体にウォッシュコートする以外は実施例1と同様にして、触媒B及びCを製造した。

実施例 4~8

実施例1で用いた酢酸銅水溶液の代わりに、 種々の金属塩水溶液を用いる外は、実施例1と 同様にして、それぞれCo, Ni, Fe, Cr, Ma でイオン交換された触媒 D, E, F, G, Hを製造 した。各触媒の製造に用いた金属塩は以下の通 りである。 する。

突施例 1

a) スラリー 脚製

バインダーとして、 8 i /A O 比が 4 0 となるように混合されたシリカソルとアルミナゾルの混合物 6 0 部に、ゼオライト (S i /A O 比 4 0; 最大細孔径 5.9 Å) 粉末 1 0 0部及び水 6 0 部を加えて充分操件し、硝酸アルミニウム裕液で p H を 3~6とし、ウォッシュコート用スラリーを調製した。

b) コーティング及び婦成

コージェライト製モノリス状ニハカム担体を水に浸漉し、余分な水を吹き払った後、上記 a)で得られたスラリーに浸漉し、取出した後余分なスラリーを圧縮空気で吹き払い、80℃で20分乾燥し、更にこれを600℃で1時間電気炉中で矯成した。

c) イオン交換

得られた焼成体を酢酸銅 (Cu (CH_aCOO)₂・ H_aO) 水溶液 (濃度 0.0 4 mo *e/_e*) を用い、

爽施例	触媒	&	萬	
4	D	酢酸コパルト	Co(CH _e	COO), 4 H, O
5	E	酢酸ニッケル	Ni (CH,	COO): 4 H:O
6	F	塩化第 二 鉄	FeCL	5 H ₂ O
7	G	硝酸クロム	Cr(NO.), • 9 H ₂ O
8	H	酢酸マンガン	Mn (CH	COO): 4 H:O

比較例

常法に従い、ァーアルミナを担体にウォッシュコートし、焼成後 Pt/Rh を担体18 当 り158/038担持した触媒 I を製造した。

試験例 1

上配各與施例及び比較例で得られた触媒 A ~ I を 3.0 & エンジンの排気系に取り付け、空燃比 (A/F) 2.0、入ガス温度 6.0 0 C の条件下 で の NOx の浄化率 を 例定した。 その結果を 第 1 表に 示す。

第 1 表

実施例系	触媒	基材(Si/Ae比)	触媒金與	NOx押化
実施例1	A	セオライト	(40)	Сu	4 5 %
奥施例 2	В	モルデライト	, (19)		3 5
実施例 5	C	フォージャサ	11(26)	•	2 0
実施例4	D	せオライト	(40)	C o	5 0
奥施例 5	E	,	(•)	Ni	3 5
実施例 6	F	,	(")	Fe	2 3
実施例 7	G	,	(•)	Сr	2 5
突施例8	Н	"	(•)	Мn	2 0
比較例	I	アーアルミナ		Pt/Rh	2

実施例9及び試験例2

パインダーのアルミナソルとシリカソルの混合比が触媒活性にどのような影響を及ぼすかをみるために、Si/A& 比が 0,50,100,200,400,100006種類のパインダーを用意し、実施例1で使用されたパインダーの代わりに上記6種類のパインダーを用い、また Cu 塩の代

イトを担体上に付着させたものであるため、リーン努囲気においてもNOxが選択的に細孔中の活性サイトに吸着・反応し、浄化される触媒となる。

従って本発明の排気ガス浄化触媒を用いれば、リーン雰囲気走行でも大気中にNOxを排出する恐れがなくなることから、エンジンの設定空燃比を大きくして、自動車の低燃費化を図ることができる。また混合気を希薄にすることでHC, CO の 発生自体も低くなる。

またバインダーのSi/AB 比をゼオライトの それに近くなるよりにすると触媒性能が向上す るとともに、パインダーとゼオライトとの体験 膨張率の急を緩和し耐制離性に富む触媒となる。

4.図面の簡単な説明

第1図は、セオライト(モルデナイト)のSi /AB比ごとのイオン交換時間とイオン交換率の 関係を示す図、

第2回は酢酸銅水溶液濃度、pHとイオン交換 塞の関係を示す図、 わりに貴金属塩を用いる以外は実施例1と同様にして各種の触媒を製造した。それらのA/F=18.0 におけるHC、CO、NOx 浄化率を調べた。その結果を第2表に示す。パインダーのSi/A&比がセオライトのSi/A&比 4 0に近い触媒は

第2表: 净化率例

Si/ABH	НC	CO	NOx
O	8 0	8 2	3 0
5 0	8 1	8 5	6 0
100	8 2	8 5	7 1
200	76	8 1	5 3
400	70	7 5	1 5
1000	5 7	6 8	1

く発明の効果>

本発明方法により得られる排気ガス浄化用触媒は、触媒能を有する金属でイオン交換されかつ NOx分子の取込みに適する細孔を持つゼオラ

第3図はゼオタイトの一例の部分構造を示す 様式図、

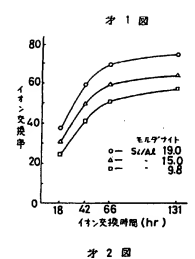
第 4 図はセオライトのイオン交換状態の説明 図である。

特 許 出 顧 人 トョタ自動車株式会社 同 株式会社 豊田 中央研究所

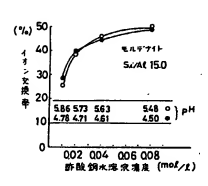
代理人 弁理士 粤 優 美 ほか2名

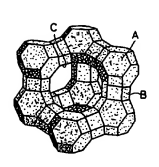


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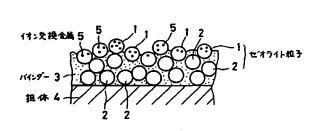


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岁 3 図



MANUFACTURING METHOD FOR AN EXHAUST GAS PURIFYING CATALYST

Masataka Kawabata et al.

UNITED STATES PATENT AND TRADEMARK OFFICE WASHINGTON, D.C. MAY 2003
TRANSLATED BY THE RALPH MCELROY TRANSLATION COMPANY

JAPANESE PATENT OFFICE PATENT JOURNAL (A)

KOKAI PATENT APPLICATION NO. HEI 1[1989]-135540

Int. Cl.⁴:

B 01 J 29/08

B 01 D 53/36

B 01 J 29/18

Sequence Nos. for Office Use.:

A-6750-4G

A-8516-4D

Filing No.:

Sho 62[1987]-292914

Filing Date:

November 19, 1987

Publication Date:

May 29, 1989

No. of Inventions:

1 (Total of 5 pages)

Examination Request:

Not filed

MANUFACTURING METHOD FOR AN EXHAUST GAS PURIFYING CATALYST

[Haiki gas jyokayo shokubai no seizo hoho]

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Applicants:

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[There are no amendments to this patent.]

Claims

- 1. A manufacturing method for an exhaust gas purifying catalyst characterized by consisting of the processes below:
 - (a) a process of mixing zeolite, silica sol, alumina sol, and water and preparing a slurry;
- (b) a process of adhering the aforementioned slurry onto an integrated type catalyst support and firing it;
- and (c) a process of ion-exchanging the zeolite by a catalytic metal through a soak in a metallic salt aqueous solution of the catalytic metal.

- 2. The method described in Claim 1 characterized by the mixing ratio between the silica sol and the alumina sol being such that the Si/Al ratio of them is not very different from the Si/Al ratio of the zeolite.
- 3. The method described in Claim 1 characterized by process (c) being a process that is performed after processes (a) and (b).

Detailed explanation of the invention

Industrial application field

This invention concerns a manufacturing method for an exhaust gas purifying catalyst for vehicles, in particular a catalyst, which can remove NOx at a high ratio even in an excess oxygen atmosphere where the air fuel ratio is on the lean side.

Prior art

As exhaust gas purifying catalysts for cars, catalysts that simultaneously attain an oxidation of carbon monoxide (CO) and hydrocarbons (HC) and a reduction of nitrogen oxides (NOx) have been widely used. With most of such catalysts, a rare metal, such as Pd, Pt, or Rh, for example, and a rare earth metal, such as Ce or La, for example, as a cocatalyst and a base metallic oxide, such as Ni, for example, in some cases are added in an alumina coated layer over a fire resisting catalyst support, as seen in Japanese Kokoku Patent No. Sho 58[1983]-20307, for example.

The purification character of such a catalyst is greatly affected by the set air fuel ratio of the engine. The amount of oxygen (O_2) is in excess after combustion in a thin air mixutre, which is on the lean side with a large air fuel ratio, which activates oxidation action and inactivates reduction action. Reversibly, oxidation action becomes inactive and reduction action becomes active on the rich side where the air fuel ratio is small. A catalyst can most effectively function near the theoretical air fuel ratio (A/F=14.6) where there is a balance between oxidation and reduction.

Accordingly, with an automobile in which an exhaust gas purifying device that uses a catalyst is attached, the oxygen concentration of the exhaust system is detected, and a feedback control is used so that the air mixture is maintained near the theoretical air fuel ratio.

Problem to be solved by the invention

On the other hand, the attainment of low fuel consumption has also been requested. For the attainment of it, it has been known that an excess oxygen air mixture should be combusted as much as possible generally during traveling. However, there is an issue, wherein when the attained air fuel ratio is an excess oxygen atmosphere on the lean side, HC and CO harmful components in the exhaust gas can be removed though an oxidation, but NOx cannot be removed through reduction because contact with an active metal is prevented by O₂, which has been adsorbed into the catalyst bed. As a result, thinning of the air mixture is not conventionally possible in automobiles that attain a high degree of purification of the exhaust gas by a catalyst.

The objective of this invention, which is made for solving the aforementioned problem, is to offer a manufacturing method for an exhaust gas purifying catalyst, which can remove NOx through reduction even at the lean side and can sufficiently eliminate all harmful components in a large region at the lean side from the theoretical air fuel ratio.

Means for solving the problem

The manufacturing method for an exhaust gas purifying catalyst in this invention has the characteristic of consisting of the processes below:

- (a) a process of mixing zeolite, silica sol, alumina sol, and water and preparing a slurry;
- (b) a process of adhering the aforementioned slurry to an integrated type carrier and firing it; and (c) a process of ion-exchanging the zeolite by a catalytic metal through soaking in a metallic salt aqueous solution of the catalytic metal.

In the process of preparing the slurry in process (a), it is ideal to mix at a proportion of 30-100 parts of a mixture of silica sol and alumina sol and 30-100 parts for water for 100 parts of zeolite. Furthermore, the obtainment of a slurry at a pH of 3-6 by adding a pH adjusting agent is desirable.

It is important for the mixing ratio between silica sol and alumina sol to be such that the Si/Al ratio of them is not very different from the Si/Al ratio of the zeolite. It is ideal to mix the silica sol with the alumina sol so that the Si/Al ratio of the mixture is 1/3-3 times the Si/Al ratio of the zeolite.

As the zeolite, zeolite having small holes at a diameter of 5-10Å, which is slightly larger than the diameter of the NOx molecule, is proper.

In process (b), monolithic and honeycomb type catalyst supports made of ceramic that are widely used are sufficient as the aforementioned integrated type catalyst support. For adhering the aforementioned slurry onto said catalyst support, the slurry may be spray-coated on the support, or the support may be immersed into the slurry. Firing is properly obtained at a temperature below 800°C after blowing away excess attached slurry by compressed air or after vacuum suction.

As catalytic metals for the ion exchange in process (c), transition metals, such as Cu, Cr, Co, Ni, Fe, and Mn, for example, and rare metals, such as Pt, Pd, Rh, Ce, Ir, and Ru, for example, can be listed. The ion exchange can be attained by soaking the zeolite into an aqueous solution of a water-soluble salt of an aforementioned metal. A proper concentration of the

solution is 0.001-0.1 mol/L. For a copper acetate aqueous solution, for example, 0.02-0.06 mol/L is particularly ideal (refer to Figure 2). A sufficient temperature of the solution is 30±10°C. The ion exchange time is 5-140 hours, ideally 40-100 hours (refer to Figure 1).

In the manufacturing method in this invention, the target catalyst can be obtained in the order of processes (a)? (b)? (c) or in the order of processes (c)? (a)? (b), however, the former order is ideal when considering the ion exchange effect because the metal that is ion-exchanged enters gradually from a site where exchange occurs easily to a more difficult site, as indicated by the structural model figure in Figure 3 in the order of A? B? C. While also considering a case, in which sites become valid? invalid for the purification of NOx in the reverse order of C? B? A, the probability for ion exchange metals 5, 5.... to enter a site higher in activity for NOx becomes higher when only ion exchanging the zeolite particles 1, 1.....that appear on the surface except for the zeolite particles 2, 2..... that are buried in the binder (3) over the catalyst support (4), as indicated in Figure 4, than when the entire zeolite is ion-exchanged at first, and the catalytic activity improves.

Operation of the invention

The catalyst that is obtained by the manufacturing method in this invention has its surface covered by zeolite, which is ion-exchanged by a catalytic metal. Zeolite has small holes in the order of several Å, which is equal to the size of the NOx molecule. As a result, NOx is selectively taken into said small holes. Sites in which the aforementioned metal is ion-exchanged and have become active are present in small holes. Therefore, NOx, which has absorbed there, is reduced by the catalytic metal.

Application examples

This invention will be explained in more detail in the application examples below.

Application Example 1

a) Preparation of the slurry

To 60 parts of a mixture of silica sol and alumina sol as a binder, which is mixed together at a Si/Al ratio of 40, 100 parts of a zeolite (Si/Al ratio of 40; maximal small hole diameter of 5.9 Å) powder and 60 parts of water are added, sufficiently stirred, the pH is adjusted to 3-6 with an aluminum nitrate solution, and a wash-coating slurry is prepared.

b) Coating and firing

A cordierite monolithic form honey-comb catalyst support is soaked in water, soaked in the slurry obtained in a) above after blowing away excess water, removed followed by excess slurry being blown away with compressed air, dried at 80°C for 20 minutes, and furthermore fired in an electric furnace at 600°C for 1 hour.

c) Ion-exchange

The obtained fired product is ion-exchanged using a copper acetate [Cu(CH₃COO)₂·H₂O] aqueous solution (concentration of 0.04 mol/L) in a penetration culturing machine for 70 hours at room temperature. After blowing away excess water, it is dried at 80°C for 20 minutes, and exhaust gas purifying catalyst A is manufactured.

Application Examples 2 and 3

Catalysts B and C are manufactured in the same manner as Application Example 1 except for using mordenite (Si/Al ratio of 19.0) and aujasite (zeolite Y; Si/Al ratio of 26) instead of the zeolite used in Application Example 1, and wash-coating slurries prepared as binder matching the respective Si/Al ratios, onto the catalyst support.

Application Examples 4-8

Catalysts D, E, F, G, and H that are ion exchanged respectively by Co, Ni, Fe, Cr, and Mn are manufactured in the same manner as Application Example 1 except for using various types of metallic salt aqueous solutions instead of the copper acetate aqueous solution used in Application Example 1. The following metallic salts were used in the manufacture of the catalysts.

突施例	触蛛	金	兵	塩	
1	D 4 mm	2 + C	CH, C	200)*	4 H ₂ O
5	B⑤ 解膜=ッ	ケル N	(CH, C	200):-	4 H ₂ O
6	F⑥ 塩化第二	二鉄 F	CL. • 6	H ₁ O	
7	G⑦ 硝酸夕 1	2 A C	r (NO.)	, • 9 H,	0
8	H⑧ 酢酸マン	ガン Mi	(CH,C	200),•	4 H ₂ O

- Key: 1 Application example
 - 2 Catalyst
 - 3 Metallic salt
 - 4 Cobalt acetate
 - 5 Nickel acetate
 - 6 Ferric chloride
 - 7 Chromium nitrate
 - 8 Manganese acetate

Comparison example

In accordance with a universal method, ?-alumina is wash-coated over a catalyst support, and catalyst I supporting 1.5g/0.3 g of Pt/Rh per 1 l of the catalyst support after firing is manufactured.

Testing example 1

Catalysts A-I obtained in the aforementioned application examples and comparison example are attached to the exhausting system of a 3.0 L engine, and the removal rate for NOx under conditions including an air fuel ratio (A/F) of 20 and a gas entering temperature of 600°C are measured. Table 1 shows the results.

Table 1

Application example No.	Catalysts	Base material (Si/Al ratio)	Catalytic metal	Purification ratio of NOx
Application No. 1	A	Zeolite (40)	Cu .	48%
Application No. 2	В	Moldelite (19)	11	55
Application No. 3	С	Forjasite (26)	11	20
Application No. 4	D	Zeolite (40)	Co	30
Application No. 5	E	" (")	Ni	35
Application No. 6	F	" (")	Ni	23
Application No. 7	G	" (")	Cr	25
Application No. 8	Н	".(")	Mn	20
Comparison Example	I	r-alumina	Pt/Rh	2

Application Example 9 and Comparison Example 2

To observe the effect of the mixing ratio between the alumina sol and silica sol in the binder on the catalytic activity, 6 types of binders with Si/Al ratios of 0, 50, 100, 200, 400, and 1000 are prepared. Various types of catalysts are manufactured in the same manner as Application Example 1 except for using the aforementioned 6 types of binders instead of the binder used in Application Example 1 and using a rare metal salt instead of the Cu salt. The HC, CO, and NOx removal rates with an A/F=18.0 are checked. Table 2 shows the results. It can be understood that catalysts having a Si/Al ratio of the binder close to the Si/Al ratio of 40 of the zeolite display an excellent purification performance.

Table 2. Removal Rate (%)

Si/Al ratio	HC	СО	NO _x
0	80	82	30
50	81	85	60
100	82	85	. 71
200	76	81	33
400	70	75	15
1000	57	68	1

Effect of the invention

The exhaust gas purifying catalyst, which is obtained by the method in this invention, has a zeolite, which is ion-exchanged by a metal having catalytic performance and which also has small holes that are suitable for taking in NOx molecules, adhered over a catalyst support, and therefore, it can become a catalyst that allows NOx to selectively adsorb and react at active sites in small holes even in a lean atmosphere and thus purifies.

Accordingly, the risk of exhausting NOx into the atmosphere during driving in a lean atmosphere is eliminated by using the exhaust gas purifying catalyst in this invention. Therefore, the set air fuel ratio of the engine can be increased, and the attainment of low fuel consumption for a vehicle can be obtained. The generation of HC and CO is also reduced by thinning the air mixture.

The catalytic performance also improves by allowing the Si/Al ratio of the binder to approach that of the zeolite, and a catalyst that is rich in separation resistance is obtained by relieving the difference in the volume expansion ratio between the binder and zeolite.

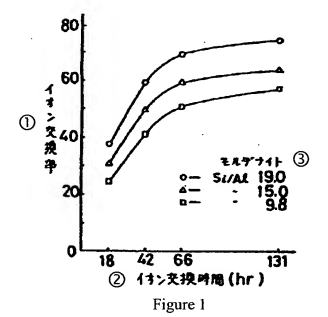
Brief description of the figures

Figure 1 is a graph showing the relationship between the ion exchange time and the ion exchange rate per the Si/Al ratio of zeolite (mordenite).

Figure 2 is a graph showing the relationship between the concentration of a copper acetate aqueous solution, pH, and the ion exchange rate.

Figure 3 is a model diagram showing a partial structure of an example of zeolite.

Figure 4 is an explanatory diagram of the ion-exchanged state of zeolite.



Keys: 1 Ion exchange rate

2 Ion exchange time (hr)

3 Mordenite

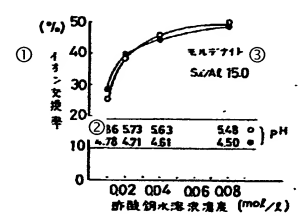


Figure 2

Keys: 1 Ion exchange rate

2 Concentration of copper acetate aqueous solution (mol/L)

3 Mordenite

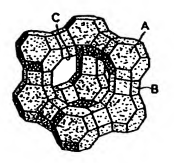


Figure 3

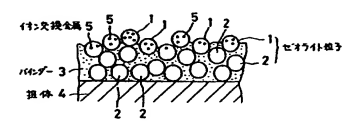


Figure 4

Keys: 1, 2

Zeolite particles Binder Catalyst support Ion exchange metal **4 5**